IN THE CLAIMS:

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Please cancel claims 1-16.

Please add new claims as follows:

- 17. (New) A method for treating heart failure in a subject, comprising:
- a) administering an angiotensin II (AT₁) receptor inhibitor to said subject for a first period beginning at about the time of a myocardial infarction;
- b) reducing administration of said angiotensin II (AT₁) receptor inhibitor after said initial period; and
- c) administering a growth hormone during a second period beginning after said reducing administration of said AT₁ receptor inhibitor.
- 18. (New) The method of claim 17, wherein said first period has a duration of about 10 to 12 weeks.
- 19. (New) The method of claim 17, wherein the AT₁ receptor inhibitor is administered at least once daily.
- 20. (New) The method of claim 17, wherein AT₁ receptor inhibitor administration is discontinued following said first period.
 - 21. (New) The method of claim 17, wherein said AT₁ receptor inhibitor comprises losartan.
- 22. (New) The method of claim 17, wherein said growth hormone is administered for about two weeks to about three months.
 - 23. (New) The method of claim 17, wherein said reducing of AT₁ receptor inhibitor allows

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for a favorable physiologic hypertrophic effect from said growth hormone.

- 24. (New) A method of treating heart failure in a subject, comprising;
- a) administering an angiotensin II (AT₁) receptor inhibitor to said subject over a first period beginning about the time of an ischemic event, and said first period continuing for a sufficient amount of time to substantially permit favorable left ventricular remodeling or limit unfavorable ventricular remodeling;
- b) decreasing said administering of AT₁ receptor inhibitor at a time approximately after said ventricular remodeling; and
- c) administering a growth hormone to said subject during a second period beginning at a time approximately after said ventricular remodeling.
- 25. (New) The method of claim 24, wherein administering said AT₁ receptor inhibitor is discontinued at about the time administering said growth hormone begins.
- 26. (New) The method of claim 24, wherein the angiotensin II (AT₁) receptor inhibitor is administered at least once daily.
- 27. (New) The method of claim 24, wherein administration of said AT₁ receptor inhibitor is discontinued at about the time administering said growth hormone begins.
- 28. (New) The method of claim 24, wherein said administration of said AT₁ receptor inhibitor following said ventricular remodeling is decreased prior to the end of said first period.
 - 29. (New) The method of claim 24, wherein said AT₁ receptor inhibitor comprises losartan.
 - 30. (New) The method of claim 24, wherein said growth hormone is human growth hormone.

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- 31. (New) The method of claim 24, wherein said AT₁ receptor inhibitor is administered beginning within seven days of said ischemic event.
- 32. (New) The method of claim 24, wherein said AT₁ receptor inhibitor is administered for about 8 to about 12 weeks.
- 33. (New) The method of claim 24, wherein said AT₁ receptor inhibitor is administered for about 10 weeks.
- 34. (New) The method of claim 24, wherein said growth hormone is administered for about two weeks to about three months.
- 35. (New) The method of claim 24, wherein a second administration of a composition comprising AT₁ receptor inhibitor is administered for a time following said growth hormone administration.
- 36. (New) The method of claim 35, wherein growth hormone is administered following said second administration of AT₁ receptor inhibitor.
- 37. (New) The method of claim 24, wherein decreasing said administering of AT₁ receptor inhibitor allows for a favorable physiologic hypertrophic effect from said growth hormone.

No new matter is introduced by these amendments.

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